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## R-E-M-SC1 SEARCH REQUEST FORM

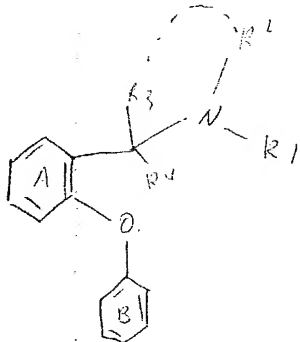
Requestor's Name: Hong Liu Serial Number: 10/075,847  
 Date: 2/9/04 Phone: 1-0669 Art Unit: 1624

**Search Topic:**

Please write a detailed statement of search topic. Describe specifically as possible the subject matter to be searched. Define any terms that may have a special meaning. Give examples or relevant citations, authors, keywords, etc., if known. For sequences, please attach a copy of the sequence. You may include a copy of the broadest and/or most relevant claim(s).

*Barb please*

FEB 12 2004  
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A, B can also be naphthyl

R<sup>2</sup>, R<sup>3</sup> can be independent, they may also form a ring.

**STAFF USE ONLY**

Date completed: 2/11/04  
 Searcher: Jayne P. Dreyer  
 Terminal time: 1:5  
 Elapsed time: 1.5 days  
 CPU time: 1.5 h  
 Total time: 1.5 h  
 Number of Searches: 1  
 Number of Databases: 3

**Search Site**

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**Vendors**

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**Type of Search**

- N.A. Sequence
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- Structure
- Bibliographic



# STIC SEARCH RESULTS FEEDBACK FORM

## Biotech-Chem Library

Questions about the scope or the results of the search? Contact *the searcher or contact:*

Mary Hale, Information Branch Supervisor  
308-4258, CM1-1E01

## Voluntary Results Feedback Form

➤ I am an examiner in Workgroup:  Example: 1610

➤ Relevant prior art found, search results used as follows:

- 102 rejection
- 103 rejection
- Cited as being of interest.
- Helped examiner better understand the invention.
- Helped examiner better understand the state of the art in their technology.

Types of relevant prior art found:

- Foreign Patent(s)
- Non-Patent Literature  
(journal articles, conference proceedings, new product announcements etc.)

➤ Relevant prior art not found:

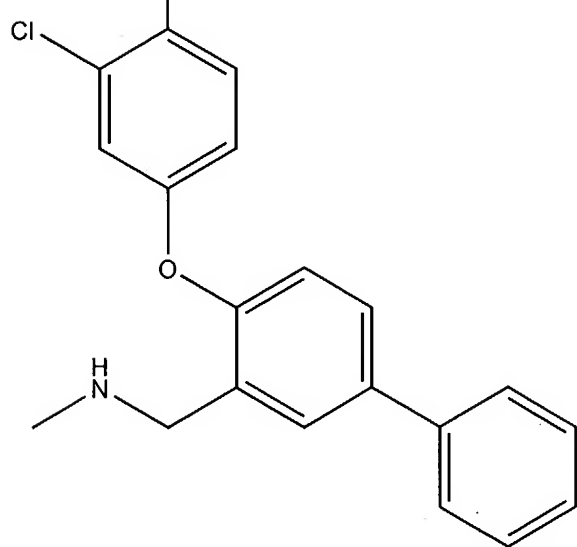
- Results verified the lack of relevant prior art (helped determine patentability).
- Results were not useful in determining patentability or understanding the invention.

Comments:

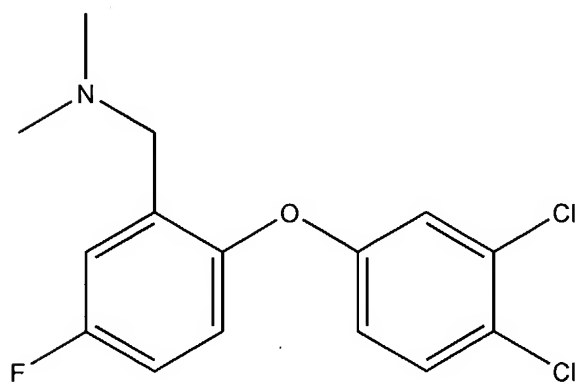
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[4-(3,4-dichlorophenoxy)-biphenyl-3-ylmethyl]-methylamine



[2-(3,4-dichlorophenoxy)-5-fluorobenzyl]-dimethylamine



**Epicupreine.** L. Prajer and J. Suszko (Poznań Univ.). *Bull. soc. amis sci. et lettres Poznań Ser. B* 13, 53-66, 67-77, 79-89 (1956) (in English).—See *C.A.* 49, 2448f.

H. M. Leicester

**Chelates and conformation of cinchona bases.** Zoltán a Földi, Tamás Földi, and András Földi (Authors' Lab., Budapest). *Chem. & Ind. (London)* 1957, 465-6.—Epiquinidine (648 mg.) ground with 5 ml. 0.2M CuSO<sub>4</sub> gives an addn. compd. which with N NaOH yields microcrystals of the chelate (C<sub>20</sub>H<sub>22</sub>O<sub>2</sub>N<sub>2</sub>)Cu. The chelate of epiquinine is similarly obtained. Both chelates decom. 150-90° and show no characteristic m.p. All attempts to prepare chelates from quinine, quinidine, cinchonine, and cinchonidine failed. The readiness to form a chelate suggests that in the epi-bases the alc. H is chelated by O and N, giving rise to a 5-membered ring and to an addnl. asymmetry absent in the C-9 epimers.

Blanche B. White

**Asymmetric induction and absolute configuration in the biphenyl series.** Jerome A. Berson and Michael A. Greenbaum (Univ. of S. California, Los Angeles). *J. Am. Chem. Soc.* 79, 2340 (1957); cf. *C.A.* 51, 1108i.—MeMgI converted the phenylglyoxylates of phenyldihydrothebaïne (I) and its derivs., 2,5,6-R'(MeO)(HO)C<sub>6</sub>H<sub>4</sub>C<sub>6</sub>H<sub>3</sub>(OME)R-3,6 (II) (IIa, R = CH<sub>2</sub>:CH<sub>2</sub>; R<sup>1</sup> = CH<sub>2</sub>CHPhNMe<sub>2</sub>) (IIb, R = Et; R<sup>1</sup> = CH<sub>2</sub>CHPhNMe<sub>2</sub>) (IIc, R = CH:CH<sub>2</sub>; R<sup>1</sup> = CH:CHPh) to atrolactic esters. Sapon. and isolation without optical fractionation gave (-)-atrolactic acid (III) (abs. configuration shown) in optical yields of 70% from I and 91, 89, and 93% from IIa, IIb, and IIc, resp. Mechanisms for the formation of III are discussed. Felix Saunders

**The structure of pseudomorphine.** K. W. Bentley and S. F. Dyk (Univ. Aberdeen, Scot.). *Chem. & Ind. (London)* 1957, 398.—Pseudomorphine was shown to be 2,2'-dimorphine (cf. Small and Turnball, *C.A.* 31, 6663<sup>a</sup>, and Goto and Kitasato, *C.A.* 24, 4299). Oxidation of 1-bromodihydromorphine with alk. K<sub>2</sub>Fe(CN)<sub>6</sub> at 70-80° gave a poor yield of dibromotetrahydropseudomorphine (I), m. above 350°, [α]<sub>D</sub><sup>25</sup> -46° ± 10° (0.518%, N HCl). It was prep'd. by bromination of tetrahydropseudomorphine in HOAc.

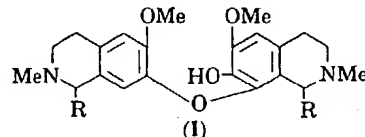
R. H. Loepert

**Synthesis in the morphinan group. II. The structure of 3-hydroxy-N-methyl-C-normorphinan.** Sciichi Saito (Univ. Tokyo). *Pharm. Bull. (Tokyo)* 4, 438-43 (1956); cf. *C.A.* 51, 8117d.—As conclusive evidence of the structure of the previously synthesized (*loc. cit.*) title compd. (I) it was submitted to the Hofmann degradation and its degradation products were synthesized. Excess CH<sub>2</sub>N<sub>2</sub> in ether added to 2.5 g. I suspended in 25 cc. MeOH, the mixt. kept 5 days at room temp., the solvents evapd., and the residue distd. *in vacuo* yielded 2.3 g. O-Me deriv. (II) of I, b<sub>10-15</sub> 152-3°; picrate, m. 167-9° (from AcOH). Refluxing 40 min. 2.8 g. MeI salt of II with 30 cc. 16% KOH, dissolving the sep'd. oil in C<sub>6</sub>H<sub>6</sub>, and distg. the residue from the C<sub>6</sub>H<sub>6</sub> soln. *in vacuo* yielded 1.6 g. 9b-(Me<sub>2</sub>NCH<sub>2</sub>CH<sub>2</sub>) deriv. (III) of 8-methoxy-2,3a,9b-tetrahydro-1H-benz[e]indene (IV), b<sub>10-06</sub> 160° (bath temp.); HCl salt, m. 181-3° (from MeOH-ether). The MeI salt of III (1.4 g.) in 30 cc. warm H<sub>2</sub>O shaken 4 hrs. at room temp. with fresh Ag<sub>2</sub>O (from 2.4 g. AgNO<sub>3</sub> and 10 cc. 3N NaOH), the filtrate evapd. to dryness below 50°, and the residue heated *in vacuo* (2-3 mm.) evolved NMe<sub>2</sub> at 90° and distd. 0.6 g. liquid at 90-140°, which, dissolved in ether, washed with 10% HCl, dried, the ether removed, and the residue distd. *in vacuo* yielded 0.5 g. 9b-(CH<sub>2</sub>:CH) deriv. (V) of IV, b<sub>1-1</sub> 130° (bath temp.). V (0.3 g.) in 20 cc. EtOH catalytically reduced (10% Pd-C) absorbed 2 molar equivs. H in 3 hrs. and yielded 0.3 g. 8-methoxy-9b-ethyl-2,3a,4,5,-9b-hexahydro-1H-benz[e]indene (VI), b<sub>1-6</sub> 100-20° (bath temp.); penta-Br deriv. (VII), m. 168-70° (decompn.);  $\lambda_{\text{CCl}_4}$  288 and 298 m $\mu$ . Synthesis of VI confirmed its structure and thus indirectly the structure of I. 2-Ethoxycarbonylcyclopentanone (VIII) (23.5 g.) added slowly to 5.6 g. K suspended in 220 cc. abs. PhMe, stirred 1 hr. at room temp., 23.5 g. *p*-MeOC<sub>6</sub>H<sub>4</sub>CH<sub>2</sub>CH<sub>2</sub>Br added dropwise, the mixt. refluxed 10 hrs., cooled, and H<sub>2</sub>O added yielded from the org. layer 20 g. 2-(*p*-MeOC<sub>6</sub>H<sub>4</sub>CH<sub>2</sub>CH<sub>2</sub>) deriv. of VIII, b<sub>1-5</sub> 168-70°, which refluxed 3 hrs. with 95 cc. AcOH, 45 cc. concd. HCl, and 150 cc. H<sub>2</sub>O, the mixt. concd. to 0.25 vol. *in vacuo*, and extd. with C<sub>6</sub>H<sub>6</sub> was converted to 12 g. 2-(*p*-methoxyphenethyl)cyclopentanone (IX), b<sub>1-6</sub> 130-7°; semi-carbazone, m. 205-7° (decompn.). IX (12 g.) in 50 cc. Ph-Me added to the Grignard reagent from 26.4 g. EtBr, 6 g. Mg, and 30 cc. abs. ether, distd. to 85°, refluxed 5 hrs., decompd. with ice H<sub>2</sub>O contg. HCl, and worked up as usual

yielded 11 g. 1-ethyl-2-(*p*-methoxyphenethyl)cyclopentene, b<sub>2</sub> 138-43°, which, added to 35 cc. 85% H<sub>2</sub>SO<sub>4</sub> at 0-5°, stirred 30 min. at 15-20°, and extd. with C<sub>6</sub>H<sub>6</sub>, was cyclized to 8.5 g. VI, b<sub>1-4</sub> 116-18°; penta-Br deriv. (X), m. 168-70° (decompn.), undepressed by VII. The ultraviolet and infrared spectra of both VI obtained by degradation and by synthesis agreed well (curves shown). Detn. of the positions of Br in X was attempted. VI (0.5 g.) in 20 cc. CHCl<sub>3</sub> kept at room temp. 1 hr. with 0.4 cc. Br, refluxed 30 min., CHCl<sub>3</sub> evapd., the residue heated 10 min. on a steam bath with 15 cc. AcOH and 0.01 cc. Br yielded 9% tetra-Br deriv. (XI) of VI, m. 172-4° (decompn.),  $\lambda_{\text{CCl}_4}$  287 m $\mu$  (log ε 3.74). X (0.7 g.) refluxed 5 hrs. with 0.7 g. MgCO<sub>3</sub>, 15 cc. dioxane, and 15 cc. H<sub>2</sub>O, poured into 30 cc. H<sub>2</sub>O, and extd. with C<sub>6</sub>H<sub>6</sub> yielded 0.35 g. (probably) 5-hydroxy deriv. (XII) of XI, m. 168-70°,  $\lambda_{\text{EtOH}}$  282 and 291 m $\mu$  (log ε 3.46 and 3.53), and this by oxidation with CrO<sub>3</sub> yielded 65% 5-oxo deriv. (XIII) of XI, m. 186-8°,  $\lambda_{\text{EtOH}}$  245 and 287 m $\mu$  (log ε 4.11 and 3.89). These results, together with the infrared spectra of X-XIII, lead to the tentative conclusion that X is the 4,5,6,7,9-Br<sub>5</sub> deriv. of VI.

H. S. French

**Alkaloid studies. XVII. The structure of the cactus alkaloid pilocereine.** Carl Djerassi, S. K. Figdor, J. M. Bobbitt, and F. X. Markley (Wayne State Univ., Detroit, Mich.). *J. Am. Chem. Soc.* 79, 2203-10 (1957); cf. *C.A.* 51, 8118d.—Structure I (R = CH<sub>2</sub>CHMe<sub>2</sub>) was elucidated for the cactus alkaloid pilocereine. I (8.5 g.) in 200 cc.



MeOH-280 cc. Et<sub>2</sub>O treated 6 days at 0° with 2.2 g. distd. CH<sub>2</sub>N<sub>2</sub>, the mixt. treated with an addnl. 2.2 g. CH<sub>2</sub>N<sub>2</sub>, kept 3 days at 0°, and evapd. and the residue recrystd. from hexane yielded 6.5 g. Me ether (II) of I, m. 92-105°, resolidified and m. 153-5° (all m.p.s. were detd. on a Köfler block). II, m. 153-5° (from EtOAc), was transformed to a 2nd cryst. form, m. 133-5°; the transformation was reversed by recrystn. from hexane. I (3.0 g.) in 100 cc. abs. EtOH treated with 3.6 g. MeCHN<sub>2</sub> in 150 cc. Et<sub>2</sub>O, kept 24 hrs. at room temp., treated with an addnl. 3.6 g. MeCHN<sub>2</sub>, refrigerated 6 days, and evapd. yielded 2.07 g. Et ether (III) of I, m. 90-5° and 152-3° (from hexane); 2nd crop, 0.32 g. Amberlite IRA-400 (HCl) (200 g.) treated with 500 cc. 50% aq. NaOH, 2 l. H<sub>2</sub>O, and finally 250 g. NaHCO<sub>3</sub> in satd. aq. soln. and washed with 12-16 l. H<sub>2</sub>O gave the bicarbonate salt IRA-400-HCO<sub>3</sub> which was stored under distd. H<sub>2</sub>O. Styphnates and picrates in EtOH or Me<sub>2</sub>CO contg. about 5% H<sub>2</sub>O passed dropwise over a column of IRA-400-HCO<sub>3</sub>, the column washed with 2 vols. 10% aq. Me<sub>2</sub>CO, the Me<sub>2</sub>CO removed *in vacuo*, acid added, the aq. soln. washed with Et<sub>2</sub>O and basified with NH<sub>4</sub>OH, and the base isolated with Et<sub>2</sub>O gave the corresponding free amines. II (2.5 g.) in 100 cc. 10% H<sub>2</sub>SO<sub>4</sub> made just alk. with 2N NaOH, treated dropwise at room temp. with 250 cc. 2% aq. KMnO<sub>4</sub>, allowed to stand overnight, acidified with H<sub>2</sub>SO<sub>4</sub>, and extd. continuously with Et<sub>2</sub>O, the residue from the ext. treated with SO<sub>2</sub>Cl<sub>2</sub> and then PhNH<sub>2</sub>, and the product chromatographed yielded 35 mg. iso-PrCONHPh and 10 mg. iso-BuCONHPh. I (5.0 g.) in 200 cc. dry Et<sub>2</sub>O added slowly with stirring to 1.5 l. liquid NH<sub>3</sub> at -60° during 5 hrs., the mixt. warmed during 3 hrs. to -30°, treated cautiously with NH<sub>4</sub>Cl and evapd. overnight, the residue partitioned between Et<sub>2</sub>O and 3% aq. NaOH, the alk. layer acidified with 40% H<sub>2</sub>SO<sub>4</sub>, washed with Et<sub>2</sub>O, basified with concd. NH<sub>4</sub>OH, and extd. with Et<sub>2</sub>O, and the ext. evapd. gave 2.46 g. phenolic basic oil (IV); the original Et<sub>2</sub>O layer extd. with 10% HCl, dried, and evapd. left only a small amt. of nonphenolic, nonbasic oil which was discarded; the acid ext. basified with NH<sub>4</sub>OH and extd. with Et<sub>2</sub>O gave 2.40 g. nonphenolic, basic, glassy material (V). V consisted mainly of *isopilocereine* (VI); dipicrate, m. 235-7° (from Me<sub>2</sub>CO). VI dipicrate (3.5 g.) treated with LiOH and the resulting free base treated with CH<sub>2</sub>N<sub>2</sub> in Et<sub>2</sub>O-MeOH yielded 55% Me ether (VII) of VI. VII 180-90° (evaporatively distd.). In 1 run, a 75-mg. i aldehyde V treated with 40 mg. picric acid yielded 70 mg. 1-isobutyl-2-methyl-6-methoxy-1,2,3,4-tetrahydroisoquinoline (VIII) picrate, m. 150-1° (from MeOH). IV (0.26 g.) treated 6 days at 0° with CH<sub>2</sub>N<sub>2</sub> in Et<sub>2</sub>O contg. a small amt.

of MeOH and evapd., the residue extd. with Et<sub>2</sub>O and washed with 3% aq. NaOH, and the resulting oil (0.2 g.) chromatographed on 9 g. Al<sub>2</sub>O<sub>3</sub> gave 0.155 g. 7-MeO deriv. (IX) of VII,  $n_D^{20}$  1.5284; *styphnate*, m. 212–13°; *picrate*, m. 184–5°. I (5 g.) in 1.5 l. dry NH<sub>3</sub> treated at –30° with 6 g. K, and the mixt. worked up in the usual manner gave 1.79 g. V and 2.68 g. IV; the IV dissolved in Et<sub>2</sub>O, dried, and concd. yielded 1.45 g. *demethylisopilocereine* (X), m. 177.5–78°. X (100 mg.) treated 2 days at 0° with excess CH<sub>2</sub>N<sub>2</sub> in Et<sub>2</sub>O and evaporatively distd. yielded 81 mg. glass, the infrared spectrum of which closely resembled that of VI; treatment with picric acid gave a small amt. of VI picrate. X (210 mg.) in Et<sub>2</sub>O-MeOH treated 7 days with CH<sub>2</sub>N<sub>2</sub> yielded 120 mg. VII. IV (300 mg.) treated 7 days at room temp. with 0.84 g. MeCHN<sub>2</sub> in Et<sub>2</sub>O, washed with alkali, and treated with picric acid gave the *picrate* of the 7-EtO deriv. of VIII, m. 151.5–2.5°. Natural IX (2.2 g.) oxidized with KMnO<sub>4</sub> yielded 310 mg. *m*-hemipinic acid, characterized as the di-Me ester, m. 89.5–90°; iso-PrCO<sub>2</sub>H and iso-Bu-CO<sub>2</sub>H were identified as their anilides. IX (2.47 g.) and 10 cc. MeI kept overnight at room temp., the resulting methiodide (5.17 g.) dissolved in a small amt. of H<sub>2</sub>O, added to 120 cc. 50% aq. KOH, and refluxed 2 hrs., and the product isolated in the usual manner yielded 2.05 g. 2,4,5-[iso-Bu(Me<sub>2</sub>N)CH](MeO)<sub>2</sub>C<sub>6</sub>H<sub>2</sub>CH:CH<sub>2</sub> (XI), oil. XI (185 mg.) in glacial AcOH ozonized 0.5 hr. at 15° and steam distd. into dimedon in MeOH, and the mixt. kept 24 hrs. at 0° gave 39 mg. CH<sub>2</sub>O deriv., m. 193–5°. XI (1.87 g.) in MeOH hydrogenated 1 hr. over 5% Pd-C yielded the 1-Et analog (XII) of XI. XII converted to the methiodide (3.94 g.) and boiled with 50% aq. KOH yielded 1.06 g. neutral N-free oil, apparently 3,4,5-Et(MeO)<sub>2</sub>C<sub>6</sub>H<sub>2</sub>CH:CHCHMe<sub>2</sub>; a 90-mg. portion ozonized and steam distd. into acidified aq. 2,4-(O<sub>2</sub>N)<sub>2</sub>C<sub>6</sub>H<sub>4</sub>NHNNH<sub>2</sub>, extd. with C<sub>6</sub>H<sub>6</sub>, and chromatographed on Al<sub>2</sub>O<sub>3</sub> yielded 20 mg. iso-PrCHO deriv., m. 181–2°. IX oxidized with KMnO<sub>4</sub> in the same manner as I gave iso-PrCO<sub>2</sub>H and iso-BuCO<sub>2</sub>H. VII (104 mg.) in C<sub>6</sub>H<sub>6</sub>—treated 4.5 hrs. with 1 cc. MeI gave 153 mg. VII.2MeI, m. 191–4° (from hexane-Me<sub>2</sub>CO). VII.2MeI (150 mg.) in 5 cc. MeOH and 20 cc. H<sub>2</sub>O passed 4 times over IRA-400-OH resin, the column washed with 20 cc. 50% aq. MeOH, and the residue from the eluates distd. yielded 89 mg. gummy methine, C<sub>33</sub>H<sub>50</sub>N<sub>2</sub>O<sub>4</sub>,  $b_{10,000}$  170–5°; a 100-mg. sample ozonized in CHCl<sub>3</sub> at –60° gave 55 mg. CH<sub>2</sub>O-dimedon deriv.; a 500-mg. sample in EtOH hydrogenated 10 min. over Pd-C yielded 450 mg. reduced methine (XIII),  $b_{10,000}$  160° (bath temp.). XIII (130 mg.) in Et<sub>2</sub>O treated with MeI, the dimethiodide (180 mg.) decompd. by the ion exchange resin method, the resulting neutral olefin (76 mg.),  $b_{10,000}$  160–80°, ozonized in CHCl<sub>3</sub> at –60°, and the distillate passed into 2,4-(O<sub>2</sub>N)<sub>2</sub>C<sub>6</sub>H<sub>4</sub>NHNNH<sub>2</sub> soln. yielded 44% 2,4-(O<sub>2</sub>N)<sub>2</sub>C<sub>6</sub>H<sub>4</sub>NHNNH<sub>2</sub>:CHCHMe<sub>2</sub> (XIIIa). II (2.56 g.) treated with MeI, the II.MeI (3.9 g.), m. 137–50° (decompn.), powd., added to 100 cc. refluxing 40% aq. NaOH, and refluxed 2.5 hrs., a 160-mg. portion of the resulting methine 4,2-, 5-R(MeO)[CH(NMe<sub>2</sub>)(CH<sub>2</sub>CHMe<sub>2</sub>)C<sub>6</sub>H<sub>2</sub>OC<sub>6</sub>H(OMe)<sub>2</sub>[CH-(NMe<sub>2</sub>)(CH<sub>2</sub>CHMe<sub>2</sub>)R-2,3,6,5 (XIV) (R = CH:CH<sub>2</sub>) (2.0 g.) ozonized in AcOH, and the mixt. steam distd. into 2,4-(O<sub>2</sub>N)<sub>2</sub>C<sub>6</sub>H<sub>4</sub>NHNNH<sub>2</sub> gave only 47 mg. CH<sub>2</sub>O deriv. XIV (R = CH:CH<sub>2</sub>) (1.9 g.) in 50 cc. 95% EtOH hydrogenated over 300 mg. 10% Pd-C, and the crude product (1.85 g.) recrystd. from MeCN gave 0.92 g. XIV (R = Et), m. 101.5–3.5°. XIV (R = Et) (1.21 g.) subjected to a 2nd stage Hofmann degradation gave 0.45 g. MeN picrate, m. 206–10°, and 0.84 g. N-free degradation product which ozonized in EtOAc at –60° and worked up in the usual manner yielded only 3% XIIIa. XIV (R = Et) converted to the dimethiodide (1.72 g.) and subjected to a Hofmann degradation in the usual manner except that the compd. was first dissolved in EtOH gave a substance,  $b_{10,000}$  155–70°, which appeared to be the di-CH(OEt)CH<sub>2</sub>CH-Me<sub>2</sub> analog (XV) of XIV (R = Et). II (1.98 g.) cleaved in the usual manner with 90 cc. Et<sub>2</sub>O, 600 cc. liquid NH<sub>3</sub>, and 2.5 g. K at –60° during 7 hrs. gave 1.30 g. nonphenolic basic and 0.67 g. phenolic basic fractions. The nonphenolic fractions dissolved in 20 cc. hexane and chromatographed on 80 g. Al<sub>2</sub>O<sub>3</sub> (deactivated with 2.4 cc. 10% AcOH), giving 114 fractions, and fractions 20–46 (hexane up to 1:1 hexane-C<sub>6</sub>H<sub>6</sub>) treated with alc. picric acid gave 0.53 g. picrate of VIII, m. 152–3°; fractions 47–83 (1:1 hexane-C<sub>6</sub>H<sub>6</sub> to 99:1 C<sub>6</sub>H<sub>6</sub>-Et<sub>2</sub>O) treated with alc. picric acid gave 0.196 g. IX picrate, m. 183–5°. Fractions 100–12 (9:1 C<sub>6</sub>H<sub>6</sub>-Et<sub>2</sub>O) gave similarly 10% picrate of the 8-OH deriv. (XVI) of IX,

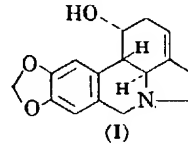
m. 150–5°. XVI (73 mg.) (from the picrate) treated 10 days at 0° with CH<sub>2</sub>N<sub>2</sub> in Et<sub>2</sub>O-MeOH and the product treated with alc. picric acid yielded the *picrate* of the 8-MeO analog (XVII) of XVI, m. 132–4°. Fractions 112–14 (Et<sub>2</sub>O and 9:1 Et<sub>2</sub>O-MeOH) gave a picrate, m. unsharply above 210°, which may represent dimeric material. The phenolic cleavage product (0.67 g.) and CH<sub>2</sub>N<sub>2</sub> in MeOH-Et<sub>2</sub>O refrigerated 8 days yielded 0.43 g. picrate of IX, m. 181–4°; the mother liquors transformed to the free amine by the ion exchange method and chromatographed on deactivated Al<sub>2</sub>O<sub>3</sub> gave 0.164 g. oil which treated with picric acid yielded 0.175 g. picrate of XVII. III (2.04 g.) in 80 cc. Et<sub>2</sub>O and 600 cc. liquid NH<sub>3</sub> treated at –60° with 3.3 g. K and the mixt. worked up after 24 hrs. gave 1.30 g. nonphenolic basic and 0.51 g. phenolic basic fractions. The nonphenolic portion chromatographed in the usual manner gave 0.576 g. VIII picrate, m. 151–3°, 0.227 g. *picrate* of the 7-EtO analog (XVIII) of IX, m. 152–3°, and 0.244 g. *picrate* of the 8-OH deriv. of XVII, m. 153–4°. The phenolic portion (0.51 g.) methylated in the usual manner and treated with picric acid gave 0.356 g. picrate of IX, m. 183–5°.

F. W. Hoffmann

Veratrum alkaloids. IV. Analysis of veratrine by paper chromatography. Karel Macek, Stanislav Vaněček, Vendula Pelcová, and Zdeněk J. Vejdělek. *Collection Czech. Chem. Communs.* 21, 1182–7 (1956) (in German).—See *C.A.* 59, 10115*i*.

E. J. C.

Alkaloids of the amaryllidaceae. X. The structure of caranine. E. W. Warnhoff and W. C. Wildman (Nat'l. Insts. of Health, Bethesda, Md.). *J. Am. Chem. Soc.* 79, 2192–8 (1957); cf. *C.A.* 50, 16803*h*; 51, 3624*d*.—A combination of degradative expts. substantiated structure



I for the alkaloid caranine. I was recovered unchanged after 2 hrs. reflux in 10% HCl, 1 hr. reflux in 10% alc. NaOH, and 4 hrs. reflux in 90% HCO<sub>2</sub>H. The pK<sub>a</sub> values were detd. in 3:7 HCONMe<sub>2</sub>-H<sub>2</sub>O for the following compds.: I 7.60,  $\alpha$ -dihydrocaranine (II) 9.00, lycorine 6.90, and dihydrolycorine 8.67. I (150 mg.) in 10 cc. dry tetrahydrofuran refluxed 25 hrs. with 150 mg. LiAlH<sub>4</sub> gave 138 mg. oily product which crystd. from EtOAc gave 111 mg. unchanged I, m. 178.5–81° (all m.p.s. are cor.). I (200 mg.) stirred 2 hrs. with 1.00 g. MnO<sub>2</sub> in 10 cc. CHCl<sub>3</sub>, filtered, and evapd., and the residual brown glass (146 mg.) sublimed at 145° and 2 mm. gave 106 mg. crude I and 40 mg. unsublimed brown residue, insol. in org. solvents and dil. HCl. I (1.00 g.) in 50 cc. H<sub>2</sub>O contg. 6 cc. 10% HCl made just basic with 10% aq. NaOH, treated with stirring with 5.0 g. KMnO<sub>4</sub> in 250 cc. H<sub>2</sub>O dropwise during 45 min., stirred 15 min., treated with SO<sub>2</sub> and then a few cc. dil. H<sub>2</sub>SO<sub>4</sub>, and extd. with EtOAc, the yellow solid residue (413 mg.) from the ext. triturated with 10% aq. KHCO<sub>3</sub> and filtered, the filtrate acidified and extd. with EtOAc, the solid residue (178 mg.) from the ext. refluxed 3.5 hrs. with 8 cc. 20% aq. NaOH under N, the mixt. acidified and extd. with EtOAc, and the residual gum (84 mg.) sublimed at 160° and 0.3 mm. gave 12.5 mg. crude *hydrastic anhydride* (III), m. 168–75°; the original aq. layer from the oxidation extd. continuously with Et<sub>2</sub>O and the resulting brown oil (83 mg.) sublimed at 160° and 0.3 mm. gave 8.5 mg. crude III, m. 140–55°. Sublimed III (4.5 mg.) recrystd. from cyclohexane-Me<sub>2</sub>CO yielded 3.0 mg. pure III, m. 179–80.5°. Crude III (8.0 mg.) triturated with 2 drops 30% aq. EtNH<sub>2</sub> and evapd., and the residue sublimed at 160° and 0.3 mm. gave 8.0 mg. *N*-ethylhydrastimide, m. 168.5–9.5° (from EtOH). I (5.000 g.) in 30 cc. glacial AcOH and 400 mg. prerduced PtO<sub>2</sub> in 5 cc. glacial AcOH hydrogenated 2 hrs., filtered, and evapd., the residue basified with 10% aq. KOH and extd. with EtOAc, and the ext. evapd. yielded 3.626 g. II, m. 170.5–72° (from EtOAc),  $[\alpha]_D^{20} -126^\circ$  (*c* 0.441) (all rotations were taken in CHCl<sub>3</sub>); *picrate*, clusters of yellow needles, m. 149–50° and 172–3° (decompn.) (from Me<sub>2</sub>CO-EtOH). II was identical with *monodeoxydihydrolycorine* (cf. Takeda and Kotera, *C.A.* 50, 16802*a*). I (300 mg.) in 9 cc. EtOH hydrogenated at ambient conditions over 100 mg. 10% Pd-C, filtered, and evapd., and the

but with morpholine, XVII, XVIII, XLI, XLII, XXIX, XXX, XXXI, XXVIII, LIII, and LIV gave 75% LXV, LXVI, 75% LXVII, LXVIII, LXII, LXIII, 80% LXIV, LXI, 80% LXIX, and LXX, resp. XI (25 g.) refluxed 15 hrs. at 50° with 16.5 g. glacial HOAc and 17 g. chloromethyl ether, H<sub>2</sub>O added, the mixt. extd. with Et<sub>2</sub>O, the Et<sub>2</sub>O layer washed, dried, and distd. gave 50% mixt. (CI), b<sub>65</sub> 140–50°, of LI and LII. CI treated with XCIX gave a mixt. of HCl salts of LIII and LIV, from which LIV, but not LIII, could be recovered by crystn. from abs. EtOH. The HCl salts of XLVI, LX, LVIII, and LV were hygroscopic and difficult to recrystallize.

J. March

**1-Oxa-7,8-benzodidehydroindolizidine from 3,4-dihydroisoquinoline and study of 1-methyl-3,4,5,6,7,8-hexahydroisoquinoline.** Woldemar Schneider and Bertold Müller (Tech. Hochschule, Karlsruhe, Ger.). *Arch. Pharm.* 294, 360–5 (1961).—3,4-Dihydroisoquinoline (I) (5 g.) in 20 ml. C<sub>6</sub>H<sub>6</sub> was treated with 4.8 g. BrCH<sub>2</sub>CH<sub>2</sub>OH 6 days at room temp. to give 92% 2-(β-hydroxyethyl)-3,4-dihydroisoquinolinium bromide (II), m. 157°, which was alkalinized with aq. NaOH to give quant. 1-oxa-7,8-benzodidehydroindolizidine (III), b<sub>65</sub> 84–7°, m. 50°, also prep'd. by treating I with ethylene oxide in MeOH 2 days. III with HBr gave II.

To 80 g. (CH<sub>2</sub>)<sub>4</sub>.CH:CCH<sub>2</sub>CH<sub>2</sub>NH<sub>2</sub> (IV) with cooling and stirring was added dropwise 60 g. AcOH, the salt heated 2 hrs. under reflux, 2 hrs. with no condenser at 160–80°, 30 min. *in vacuo* at 120° to remove H<sub>2</sub>O, and then distd. to give 89% the Ac deriv. (V), m. 53°. V (70 g.) in 350 ml. C<sub>6</sub>H<sub>6</sub> refluxed 3 hrs. with 75 g. POCl<sub>3</sub> gave 42% 1-methyl-5,6,7,8-tetrahydroisoquinoline [b<sub>12</sub> 114–16°; HCl salt m. 233° (decompn.); HBr salt m. 231°], and 35% 1-methyl-1,2,3,4,5,6,7,8-octahydroisoquinoline, b<sub>12</sub> 101–2° (*N*-p-nitrobenzoyl deriv. m. 101°; 9,10-dibromide HBr salt m. 145–7° (decompn.); NAc deriv. b<sub>12</sub> 158°). The *N*-formyl deriv. of IV failed to cyclize with POCl<sub>3</sub> to hexahydroisoquinoline.

Norman Hosansky

**Synthesis and halomethylation of bis(3,4-dimethoxyphenyl) ether. Reaction of halomethyl derivatives with secondary amines and pyridines.** Élisabeth Matarasso-Tchiroukhine (Sorbonne, Paris). *Compt. rend.* 250, 1867–9 (1960).—[3,4-(MeO)<sub>2</sub>C<sub>6</sub>H<sub>3</sub>]<sub>2</sub>O (I) was prep'd. by refluxing 3,4-(MeO)<sub>2</sub>C<sub>6</sub>H<sub>3</sub>OK, 3,4-(MeO)<sub>2</sub>C<sub>6</sub>H<sub>3</sub>I, and Cu powder in HCONMe<sub>2</sub> 18 hrs. Ether extrn. of the dild., acidified mixt. and removal of the ether gave I, m. 94.5–95°. Treatment of I with MeCl in glacial HOAc gave [2,4,5-Cl(MeO)<sub>2</sub>C<sub>6</sub>H<sub>3</sub>]<sub>2</sub>O (II), m. 121–22°. I, MeCl, HI, and Ac<sub>2</sub>O gave the 2-ICH<sub>2</sub> analog (III) of II, m. 152°. The following derivs. of II and III were prep'd.: pyridinium salts of II and III, m.p. not given and m. 172–3° (decompn.) (MeOH), resp.; isoquinolinium salts of II and III, m. 188–9° (decompn.) (MeOH–HCONMe<sub>2</sub>) and m. 203–5° (decompn.) (MeOH), resp.; 2-morpholinomethyl analog of II, m. 216–17° (EtOH); and the 2-(Et<sub>2</sub>NCH<sub>2</sub>) analog of II, m. 142–4° (EtOH).

John W. Hylin

**The synthesis of esters of some amino acids having pharmacological importance. I. The synthesis of esters of piperidino carboxylic acids.** Béla Matkovics, Sándor Foldeak, János Pórsász, and György Sipos (Tudományegyetem, Szeged, Hung.). *Acta Pharm. Hung.* 31, 113–21 (1961) (in Hungarian).—RCH<sub>2</sub>CO<sub>2</sub>R' (I), RCH<sub>2</sub>CH<sub>2</sub>CO<sub>2</sub>R' (II), BzOCH<sub>2</sub>CH<sub>2</sub>R (III), and AcOCHMeCH<sub>2</sub>R (IV) were prep'd. I were prep'd. by condensing CICH<sub>2</sub>CO<sub>2</sub>R' with a secondary amine, II by boiling CICH<sub>2</sub>CH<sub>2</sub>CO<sub>2</sub>R' with the amine, and III by the reaction of an amino alc. with BzCl. The following I were obtained (R, R', b.p./mm., m.p. of picrate, m.p. of HCl salt, and m.p. of methiodide are given): piperidino, Me, 69°/5, 115°, 214°, 163–4°; piperidino, Et, 68°/1, 122°, 117–17.5°, 160–60.3°; piperidino, Bu, 100–1°/4, 85°, —, 178°; piperidino, PhCH<sub>3</sub>, 134–5°/1, 137°, 133°, 91–6°; morpholino, Me, 77°/2, 143°, 150.5°, 147.5°; morpholino, Et, 86–7°/4, 163°, 181°, 132–3°; morpholino, Bu, 105.5–106°/3, —, 127–9°, 95–6°; morpholino, PhCH<sub>3</sub>, 184–5°/5, 143°, 149°, —; pyrrolidino, Me, 72–3°/8, 104°, —, 153°; pyrrolidino, Et, 59–60°/2, 119.5°, 133–3.5°, —; pyrrolidino, Bu, 81–2°/3, 109.5°, —, —; pyrrolidino, PhCH<sub>3</sub>, 134–5°/1, 159–60°, 139–40°, 156°. The following II were prep'd. (data as above): piperidino, Me, 72°/2, 164°, 189°, 147–8°; piperidino, Et, 102–3°/5, 131.5°, 169°, —; piperidino, Bu, 124–5°/6, 108–9°, 164.7°, —; piperidino, PhCH<sub>3</sub>, 149–50°/1, 113°, 193.5°, —; piperidino, Ph, 114–20°/3, —, 192–5°, —; piperidino, CPh, 171°/1, —, 214°, —; morpholino, Me, 82°/2, 129°, 203°, 151°; mor-

pholino, Et, 108°/6, 108°, 188–9°, —; morpholino, Bu, 131–2°/6, 150°, 173°, 115°; morpholino, PhCH<sub>3</sub>, 154°/1, 125°, 189–90°, —; pyrrolidino, Me, 76°/5, 147°, 123°, 166°; pyrrolidino, Et, 85°/6, 114°, 146°, —; pyrrolidino, Bu, 106–8°/5, 97°, 74–5°, 115°; pyrrolidino, PhCH<sub>2</sub>, 145–6°/3, 102°, 152°, 154°. IV (R = pyrrolidino) (V), b<sub>5</sub> 75°, picrate m. 111–12°, gave a hygroscopic HCl salt. III (R = piperidino) b<sub>2</sub> 141°; HCl salt m. 184°; methiodide m. 141.5°. The action of the compds. on blood pressure and on respiration was given. II (R = N-piperidino, R' = CPh) and V had strong antinicotinic action. The effect of the piperidino and pyrrolidino propionates was increased by quaternization.

E. Kasztreiner

**Molecular structure of cyclic compounds containing sulfur.** Kenjiro Hayasaki (Tokyo Gakugei Univ.). *J. Sci. Hiroshima Univ. Ser. A* 24, 679–90 (1960).—Dipole moments of tri(thiobenzaldehyde) (I) (Baumann and Fromm, *Ber.*, 24, 1436 (1891)), tri(thio-*p*-bromo- (II), and *p*-chlorobenzaldehyde (III) were detd. The values were used to investigate the structures and isomerism of trithiane rings. Infrared and Raman spectra of 1,4-dithiane (IV) were measured, and the structure of this ring system discussed. The skeletal frequencies of IV were calcd. by Wilson's method, assuming the Urey-Bradley-Shimanouchi field. III was prep'd. by the Wörner method (*Ber.* 29, 154 (1896));  $\alpha$ -isomer m. 162°,  $\beta$ -isomer 195.5°. Dipole moments reported were (compd., moment for  $\alpha$ - and  $\beta$ -isomer given in D.): I, 2.09, 2.08; II, 2.17, 3.70; III, 2.21, 3.67. The  $\alpha$ -isomers of I–III were assigned the chair (*a,e,e*) configuration and the  $\beta$ -isomers the chair (*e,e,e*) configuration. IV also had a chair configuration.

H. H. Jaffé

**Preparation and polymerization of *S,S'*-divinyldithiocarbonate.** Helmut Ringsdorf and C. G. Overberger (Polytech. Inst. of Brooklyn, Brooklyn, N.Y.). *Makromol. Chem.* 44–46, 418–26 (1961).—The title compd. (I) in benzene with free radical initiation gave a sol. polymer contg. the structural unit SC(O)SCH(CH<sub>2</sub>)—CH<sub>2</sub>CH— and some

residual unsatn. A soln. of 20 g. ethylene sulfide (II), 51 g. COCl<sub>2</sub> (III), and 3 drops pyridine was stirred 2 hrs. at –10 to –5°, then kept 10 hrs. at 25°, the excess III removed in a stream of N, and the residue distd. to give 65% CICOSCH<sub>2</sub>CH<sub>2</sub>Cl (IV), b<sub>5</sub> 57.5°, a strong lacrimator and vesicant. A mixt. of 15.9 g. IV in 100 cc. CHCl<sub>3</sub> and 31.2 g. BrCH<sub>2</sub>CH<sub>2</sub>NH<sub>2</sub>·HBr vigorously stirred at 0° with 100 cc. 8% NaOH gave 91% BrCH<sub>2</sub>CH<sub>2</sub>NHCOSCH<sub>2</sub>CH<sub>2</sub>Cl, m. 89–90°. IV and PhSNa gave 93% PhSCOSCH<sub>2</sub>CH<sub>2</sub>Cl, b<sub>5</sub> 114–15°. II (60 g.), 49.6 g. III, and 3 drops pyridine kept at –5° and then 10 hrs. at 60° gave (on distn.) 12 g. IV and 76% CO(SCH<sub>2</sub>CH<sub>2</sub>Cl), (V), b<sub>5</sub> 96–7°, m. 40–1°, a vesicant. V (60 g.) in 150 cc. anhyd. *tert*-BuOH was added dropwise to 60.5 g. *tert*-BuOK in 405 ml. *tert*-BuOH while the mixt. warmed to 50°. The mixt. was boiled 3 hrs., neutralized with HOAc, filtered, and distd. Redistn. of the fraction b<sub>5</sub> 60–80° gave 11% I, b<sub>5</sub> 73–4°. Other products of this reaction were (*tert*-Bu)<sub>2</sub>CO, CH<sub>2</sub>:CHSCO<sub>2</sub>Bu-*tert*, and II. Polymerization of I was initiated by (NCCMe<sub>2</sub>N<sub>2</sub>)<sub>2</sub> (VI). In bulk, conversion of 20% or more gave insol. polymers swelled by C<sub>6</sub>H<sub>6</sub>, CHCl<sub>3</sub>, and HCONMe<sub>2</sub>. C<sub>6</sub>H<sub>6</sub> solns. contg. approx. 1–30% I and 0.7–1.5% VI (calcd. on I) were polymerized at 60° under N with 11.5–60.9% conversion. The polymers were filtered, and pptd. from CHCl<sub>3</sub> soln. by MeOH. They softened at 300–10° with discoloration from 280° and rapid decompn. above 310°. The infrared spectrum showed only very weak absorption at 1590 cm<sup>–1</sup> (vinyl group). Hydrolysis with KOH–MeOH under N gave a (CH<sub>2</sub>CHSH)<sub>n</sub> sol. in dil. NaOH, cross-linked by traces of O.

Otto S. Kauder

**Condensation of ethyl nitroacetate with *o*-aminophenyl mercaptan.** A. I. Kipriyanov and T. M. Verbovskaya (Inst. Org. Chem., Kiev). *Zhur. Obshchel Khim.* 31, 531–7 (1961); cf. CA 50, 9387c; Mills, CA 16, 1954.—Heating *o*-H,N<sub>2</sub>C<sub>6</sub>H<sub>4</sub>SH with O<sub>2</sub>NCH<sub>2</sub>CO<sub>2</sub>Et 4 hrs. at 100° gave 74% 2,3-dioxodihydrobenzo-1,4-thiazine 2-oxime, decompd. at 267°, also formed from HONH<sub>2</sub> and 2,2-dichloro-3-oxido-dihydrobenzo-1,4-thiazine (I) in EtOH in 79% yield; the oxime formed a mono-K salt, yellow, decompd. at 270°. The latter heated in xylene with Me<sub>2</sub>SO<sub>4</sub> 6 hrs. gave 79% Me ether (II), m. 251°, also formed from I and MeONH<sub>2</sub> in EtOH. The oxime refluxed with Ac<sub>2</sub>O 2 hrs. gave the monoacetate, decompd. at 218°; BzCl in pyridine similarly gave monobenzoate, decompd. at 235°. Heating *o*-MeNH-C<sub>6</sub>H<sub>4</sub>SH with O<sub>2</sub>NCH<sub>2</sub>CO<sub>2</sub>Et 2 hrs. at 100° gave 54% 2,3-

2,4- and 3,5-Et<sub>2</sub>C<sub>6</sub>H<sub>3</sub>OH. *o*-EtC<sub>6</sub>H<sub>3</sub>OH b<sub>765.7</sub> 202–3°, d<sub>20</sub> 1.0177, n<sub>D</sub><sup>20</sup> 1.5363; *p*-isomer b<sub>765.7</sub> 214–15°, 1.0097, 1.5328; 2,4-Et<sub>2</sub>C<sub>6</sub>H<sub>3</sub>OH b<sub>765.7</sub> 229–30°, 0.9811, 1.5264; 3,5-isomer b<sub>765.7</sub> 242.5–5°, m. 76–6.5°. The neutral oil contains EtOPh and EtC<sub>6</sub>H<sub>3</sub>OEt.

G. M. Kosolapoff

Synthesis of 2,6-diisopropylphenol. Shigeru Tsutsumi, Tadashi Yoshizawa, and Kikuhiko Koyama (Osaka Univ.). *Nippon Kagaku Zasshi* 77, 737–8 (1956).—In C.A. 52, 304b, lines 8 and 9, the compd. should be 2-isopropyl-4-chlorophenol, b<sub>7</sub> 121–26°, and I should be 2,6-diisopropyl-4-chlorophenol.

R. E. S.

Triphenyloxonium salts. A. N. Nesmeyanov and T. P. Tolstaya (M. V. Lomonosov State Univ., Moscow). *Doklady Akad. Nauk S.S.R.* 117, 626–8 (1957).—Ph<sub>3</sub>O (150 g.) treated at 80–90° with 10.5 g. PhN<sub>2</sub>BF<sub>4</sub> in 300 ml. Me<sub>2</sub>CO, heated 0.5 hr., cooled, washed with 50% Me<sub>2</sub>CO, the filtrate extd. with Et<sub>2</sub>O, and the ext. evapd. gave 2% Ph<sub>3</sub>OB<sub>4</sub>, decomp. 226° (Et<sub>2</sub>O–Me<sub>2</sub>CO). Also prep'd. were: 62% Ph<sub>3</sub>OCl, decomp. 193–4°; 72% Ph<sub>3</sub>OB<sub>4</sub>, decomp. 182–2.5°; Ph<sub>3</sub>OI, decomp. 177–8°; Ph<sub>3</sub>OHgI<sub>2</sub>, decomp. 156–7°; Ph<sub>3</sub>OBPh<sub>4</sub>, decomp. about 165°; Ph<sub>3</sub>OPtCl<sub>4</sub>, decomp. 184–5°; Ph<sub>3</sub>OCr<sub>2</sub>O, decomp. 180°; Ph<sub>3</sub>OICl<sub>4</sub>, decomp. 167–71°; Ph<sub>3</sub>O picrate, decomp. 155–7°. Refluxing Ph<sub>3</sub>OB<sub>4</sub> 25 hrs. in H<sub>2</sub>O left some 50% unchanged. Such refluxing with aq. NaNO<sub>2</sub> gave some 25% PhNO<sub>2</sub> isolated after reduction to PhNH<sub>2</sub>. Refluxing Ph<sub>3</sub>OB<sub>4</sub> with aq. NaN<sub>3</sub> 14.5 hrs. gave 27% PhN<sub>3</sub>, isolated after reduction to PhNH<sub>2</sub>. Ph<sub>3</sub>OB<sub>4</sub> refluxed 8.5 hrs. with aq. Et<sub>2</sub>NH gave 59% PhNET<sub>2</sub>, isolated by azo coupling with nitraniline. Refluxing Ph<sub>3</sub>OB<sub>4</sub> in pyridine 4 hrs. gave 1-phenylpyridinium fluoroborate, 89%, m. 177.5–8.5°. Absorption spectra of the Ph<sub>3</sub>O salts are reproduced.

G. M. Kosolapoff

Preparation and transformation of *p*-diethylbenzene hydroperoxide. P. G. Sergeev and A. M. Sladkov. *Zhur. Obrabch. Khim.* 27, 3349–53 (1957).—Reduction of *p*-AcC<sub>6</sub>H<sub>4</sub>Et with 80% NaH<sub>2</sub>O and KOH in O(CH<sub>2</sub>CH<sub>2</sub>OH), at 260° gave about 20% *p*-C<sub>6</sub>H<sub>4</sub>Et<sub>2</sub>, b<sub>7</sub> 70°, d<sub>20</sub> 0.860, n<sub>D</sub><sup>20</sup> 1.4947. This percolated with air at 110° in the presence of Ni(OBz)<sub>2</sub> 15–18 hrs. gave after treatment with aq. NaOH and extn. with Et<sub>2</sub>O followed by percolation of the alk. soln. with CO<sub>2</sub> and extn. with Et<sub>2</sub>O an unstated yield (about 16%) of *p*-diethylbenzene hydroperoxide, n<sub>D</sub><sup>20</sup> 1.5231, 94.2% assay. This heated in Me<sub>2</sub>CPh 3 hrs. at 130° gave 75% *p*-AcC<sub>6</sub>H<sub>4</sub>Et. Reduction of the hydroperoxide with LiAlH<sub>4</sub> gave 70% *p*-EtC<sub>6</sub>H<sub>4</sub>CHMeOH, b<sub>7</sub> 121–2°. Stirring the hydroperoxide in C<sub>6</sub>H<sub>6</sub> with 1 drop H<sub>2</sub>SO<sub>4</sub> 1 hr. gave C<sub>6</sub>H<sub>6</sub>, *p*-EtC<sub>6</sub>H<sub>4</sub>OH, and AcH.

G. M. Kosolapoff

The rearrangement of benzenesulfonyl chloride with sodium iodide in acetone. H. Kroepelin and K. Born (Tech. Hochschule, Braunschweig, Ger.). *Arch. Pharm.* 287, 561–5 (1954); cf. C.A. 21, 573.—Treating 15 g. f PhSO<sub>2</sub>Cl with 26 g. NaI in 200 ml. Me<sub>2</sub>CO gives after 2½ hrs. and subsequent processing 60.3% Na benzenesulfinate, 27.8% diphenyl disulfone, m. 191–2°, and 10.5% Ph phenylthiosulfinate, m. 41–2°. The reaction mechanism is discussed.

Henry B. Hastie

Some new phenethylamines. J. R. Merchant and A. J. Mountvala (Inst. Sci., Bombay). *Current Sci. (India)* 26, 211–12 (1957).—A series of phenethylamines was prep'd. by the reaction of an aldehyde and MeNO<sub>2</sub> in the presence of AcOH and NH<sub>4</sub>OAc to give a  $\beta$ -nitrostyrene which was then reduced with LiAlH<sub>4</sub>. The following substituted phenethylamines were isolated as their picrates (substituents and m.p. given): 2,4,6-(MeO)<sub>2</sub>Me, 117°; 2,4,6-(EtO)<sub>2</sub>Me, 115°; 2,4,6-(EtO)(MeO)Me, 135°; 2,6,4-Me<sub>2</sub>(MeO), 115°; 2,6,4-Me<sub>2</sub>(EtO), 81°; 2,4,6-Me<sub>2</sub>(MeO), 140°; 2,4,6-Me<sub>2</sub>(EtO), 113°; 2,3-PhCH<sub>2</sub>O(MeO), —(oil); and 2,3,5-(MeO)<sub>3</sub>, 102°.

P. McElroy

New method of syntheses of musk ambrette. Hiroshi Horiguchi (Kobe Univ.). *Koryo* No. 47, 18–39 (1957).—Musk ambrette (I) was synthesized from *o*-nitrotoluene. Thus, *o*-nitrotoluene was reduced to *o*-tolylhydroxylamine (II) with Zn powder in MeOH. When II was heated in MeOH with concd. H<sub>2</sub>SO<sub>4</sub>, II was rearranged to amino-*m*-cresol Me ether (III). *m*-Cresol Me ether (IV) was prep'd. by a deamination of diazotized III. The yield of IV was 45% from *o*-nitrotoluene. I was prep'd. from IV as usual.

S. Inokawa

Synthesis of guanidine compounds of diphenyl ether. I. Genzo Ito (Pharm. Hochschule Meiji, Tokyo). *Pharm. Bull. (Tokyo)* 5, 397–400 (1957) (in German).—To find new compds. active against tuberculosis, there were synthesized 8 derivs. of PhOC<sub>6</sub>H<sub>4</sub>NHC(:NH)NH<sub>2</sub> (I) and 3 derivs. of

PhOC<sub>6</sub>H<sub>4</sub>CH<sub>2</sub>NHC(:NH)NH<sub>2</sub> (II). The HCl salt of 4-H<sub>2</sub>NC<sub>6</sub>H<sub>4</sub>OPh (III) (5.5 g.) and 1.6 g. H<sub>2</sub>NCO refluxed 3 hrs. in 35 cc. abs. EtOH, the solvent removed, and the sirupy residue dissolved in H<sub>2</sub>O and made alk. with NaOH yielded 5.9

a g. I (guanidino group in 4-position), m. 137° (C<sub>6</sub>H<sub>6</sub>); nitrate, m. 173°; picrate, m. 205°. Similarly from 2- and 3-H<sub>2</sub>N derivs. of Ph<sub>2</sub>O were synthesized I (guanidino group in 2- and 3-positions), nitrates, m. 139° and 173°, resp.; and from the 4'-Me, 4'-Cl, 4'-Br, and 4'-guanidino derivs. of III, the corresponding derivs. of I, nitrates, m. 162°, 186°, 189°, and 223°, resp. The 4'-HO deriv. of I was prep'd. in 3

steps: adding 2.5 g. 4,4'-O<sub>2</sub>N(MeO) deriv. (IV) of Ph<sub>2</sub>O to 2.7 g. dry AlCl<sub>3</sub> in 20 cc. warm PhNO<sub>2</sub>, heating the mixt. 2 hrs. at 50–5°, pouring it gradually into H<sub>2</sub>O contg. 10 cc. concd. HCl and ice, steam-distg. the org. layer to remove PhNO<sub>2</sub>, and extg. the residue with hot 5% NaOH yielded from the ext. 1 g. solid unchanged IV, and from the acidified filtrate 1 g. 4,4'-O<sub>2</sub>N(HO) deriv. (V) of Ph<sub>2</sub>O, m. 172° (C<sub>6</sub>H<sub>6</sub>). V (2.3 g.) reduced in 20 cc. abs. EtOH with 3 g.

N<sub>2</sub>H<sub>4</sub>·H<sub>2</sub>O and a little Raney Ni (Balcom and Furst, C.A. 49, 8158d) yielded 1.8 g. of the corresponding 4,4'-H<sub>2</sub>N(HO) deriv., m. 152° (C<sub>6</sub>H<sub>6</sub>), and this treated as III was with H<sub>2</sub>N-CN gave the 4'-HO deriv. of I, m. 223°; HCl salt, m. 242°; flavianate, m. 200°. For the prepn. of II, 92 g. 4-MeC<sub>6</sub>H<sub>4</sub>OPh in 200 g. (CH<sub>2</sub>Br)<sub>2</sub> gently boiling on an oil bath was treated dropwise in sunlight with 80 g. Br in 50 g. (CH<sub>2</sub>Br)<sub>2</sub>, with stirring during 1 hr., stirred an addnl. 1 hr., and the cooled mixt. neutralized with solid K<sub>2</sub>CO<sub>3</sub> and distd. *in vacuo* to yield 97 g. 4-BrCH<sub>2</sub>C<sub>6</sub>H<sub>4</sub>OPh (VI), b<sub>6</sub> 157–60°. 2-Br-CH<sub>2</sub>C<sub>6</sub>H<sub>4</sub>OPh, b<sub>6</sub> 135–40°, was similarly prep'd. Dry HCl passed through 68 g. VI, 38 g. (CH<sub>2</sub>)<sub>6</sub>N<sub>4</sub>, and 42 g. NaI in 350 cc. 95% EtOH ppts. NH<sub>4</sub>Cl, and the filtrate evapd.

d yielded 25 g. HCl salt of 4-H<sub>2</sub>NCH<sub>2</sub>C<sub>6</sub>H<sub>4</sub>OPh (VII), which in concd. aq. soln. made alk. and extd. with CHCl<sub>3</sub> gave an oil; nitrate, m. 170° (decompn.) (H<sub>2</sub>O); benzoate, m. 127° (MeOH). 2-H<sub>2</sub>NCH<sub>2</sub>C<sub>6</sub>H<sub>4</sub>OPh (VIII), nitrate, m. 153° (decompn.) (H<sub>2</sub>O), was similarly prep'd. Similarly, from 26.4 g. 4-O<sub>2</sub>N deriv. of the Cl analog of VI (Southwick, et al., C.A. 49, 955c) was prep'd. 10 g. HCl salt of the 4'-O<sub>2</sub>N deriv. of VII; the free base an orange-yellow viscous oil; Ac deriv. (IX), m. 120°. IX (28.6 g.) reduced as V was yielded 21 g. 4-AcNHCH<sub>2</sub>C<sub>6</sub>H<sub>4</sub>OC<sub>6</sub>H<sub>4</sub>NH<sub>2</sub>-4' (X), m. 146°. X (12.8 g.) diazotized in the usual way, the mixt. refluxed 3 hrs. with 2 g. urea, and worked up as usual yielded 3.1 g. 4'-HO deriv. (XI) of VII, m. 182° (EtOH); picrolonate, m. 230–2° (decompn.) (MeOH-AcOEt). VII (2 g.) in 30 cc. abs. EtOH refluxed 3 hrs. with 2.6 g. MeSC(:NH)NH<sub>2</sub>·HI (XII), EtOH distd. off, and the residue in a little H<sub>2</sub>O treated with NH<sub>4</sub>NO<sub>3</sub> gave the nitrate of II, m. 157° (Me<sub>2</sub>CO-AcOEt).

Similar treatment of VIII gave the nitrate of its corresponding guanidino compd., m. 132°. XI (1.8 g.) in 20 cc. abs. EtOH refluxed 2 hrs. with 2.2 g. XII yielded on evapn. of EtOH 2.4 g. 4'-HO deriv. of II; HI salt, m. 227° (decompn.); picrolonate, m. 265–70° (decompn.) (EtOH). All 11 guanidine derivs. gave a pos. Sakaguchi reaction, a violet-red color with 2-naphthol and NaOBr. The effect of these 11 compds. against tuberculosis bacilli *in vitro* was detd. according to Tomita and Watanabe (C.A. 46, 7617h), and none was very effective. II. A new synthesis of diphenyl ether aldehyde by the Sommelet reaction and experiments with methylguanidine derivatives. 1. *Ibid.* 401–5.

Four compds. similar to the preceding but contg. 2 CH<sub>2</sub> groups between the guanidino group and the Ph<sub>2</sub>O nucleus were prep'd. According to the Sommelet reaction 66 g. 4-BrCH<sub>2</sub>C<sub>6</sub>H<sub>4</sub>OPh (I) and 38.5 g. (CH<sub>2</sub>)<sub>6</sub>N<sub>4</sub> in 200 cc. CHCl<sub>3</sub> were refluxed 4 hrs., and cooled to yield 85 g. of the condensation product, which (101 g.) with 35 g. addnl. (CH<sub>2</sub>)<sub>6</sub>N<sub>4</sub> was hydrolyzed by refluxing 2 hrs. with 400 cc. 50% AcOH, an addnl. 10 min. with 100 cc. concd. HCl, cooling, and extg. with ether to yield 33.5 g. 4-OHC-C<sub>6</sub>H<sub>4</sub>OPh (II), b<sub>6</sub> 157–60°; phenylhydrazone, m. 144°; semicarbazone, m. 214–15°. 2-OHCC<sub>6</sub>H<sub>4</sub>OPh (III) was similarly prep'd., b<sub>6</sub> 156–9; semicarbazone, m. 207–8°. Bromination of 107 g. 4'-MeO deriv. of 4-MeC<sub>6</sub>H<sub>4</sub>OPh with Br in (CH<sub>2</sub>Br)<sub>2</sub> (preceding abstr.) yielded the 4'-MeO deriv. of I, which without isolation underwent the Sommelet reaction (like I above) to yield 51 g. 4'-MeO deriv. (IV) of II, m. 57–9° (petr. ether); semicarbazone, m. 212°. Reaction

with MeNO<sub>2</sub> changed the CHO group of II, III, and IV to O<sub>2</sub>NCH<sub>2</sub>CH (IIa, IIIa, and IVa); IIa, m. 100°, 21 g. from 30 g. II; IIIa, m. 107°; IVa, m. 77°, 11.3 g. from 38 g. IV. IIa, IIIa, and IVa were reduced by LiAlH<sub>4</sub> in the usual way to the corresponding H<sub>2</sub>NCH<sub>2</sub>CH<sub>2</sub> derivs. (IIb, IIIb, IVb):

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